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510(k) Summary Information:

Device Manufacturer: Dade Behring Inc.

Contact name:

Maureen Mende, Regulatory Affairs Group Manager

Phone/Fax:

916-374-3174/916-374-3144

Date prepared:

August 28, 2006

Product Name:

Microdilution Minimum Inhibitory Concentration (MIC) Panels

Trade Name:

MicroScan MICroSTREP plus® Panel

Intended Use:

To determine bacterial susceptibility to Clindamycin For determining antimicrobic susceptibility with aerobic

streptococci including Streptococcus pneumoniae

Predicate device:

Indication for Use:

MicroScan® MICroSTREP plus® Panel

510(k) Summary:

The MicroScan MICroSTREP *plus*[®] Panel is used to determine quantitative and/or qualitative antimicrobial agent susceptibility of colonies grown on solid media of aerobic streptococci, including *Streptococcus pneumoniae*. After inoculation, panels are incubated for 20 – 24 hours at 35°C +/- 1°C in a non-CO2 incubator, and read according to the Package Insert.

The antimicrobial susceptibility tests are miniaturizations of the broth dilution susceptibility test. Various antimicrobial agents are diluted in water, buffer or minute concentrations of broth to concentrations bridging the range of clinical interest. Panels are rehydrated with 115 µl Mueller-Hinton broth supplemented with 2-5% lysed horse blood (LHB) and buffered with 50 mM HEPES, after inoculation of the broth with a standardized suspension of the organism in saline. After incubation in a non-CO2 incubator for 20-24 hours, the minimum inhibitory concentration (MIC) for the test organism is manually read by observing the lowest antimicrobial concentration showing inhibition of growth. Additionally, the panels may be incubated in and read by a MicroScan® WalkAway instrument.

The proposed instrument read method for the MicroScan MICroSTREP *plus®* Panel demonstrated substantially equivalent performance with streptococcal isolates when compared with an expected result generated on a CLSI frozen Reference Panel, as defined in the FDA document "Class II Special Controls Guidance Document: Antimicrobial Susceptibility Test (AST) Systems; Guidance for Industry and FDA", dated February 5, 2003.

This Premarket Notification (510[k]) presents data in support of reading the MICroSTREP plus[®] Panel with Clindamycin on the MicroScan[®] WalkAway instrument.

The external evaluations conducted with CDC Challenge strains were designed to confirm the acceptability of the proposed instrument read method for the MICroSTREP *plus*[®] Panel by comparing its performance with Expected Results determined before the evaluation.

In addition, data on fresh and stock isolates is presented from the original external design validation of the manual method to support removing the "Do not report" limitation for *S. pneumoniae* with Clindamycin.

The MICroSTREP *plus*® Panel demonstrated acceptable performance with an overall Essential Agreement of \geq 95.7% for Clindamycin compared with the Expected Result including *S. pneumoniae*.

Instrument reproducibility testing demonstrated acceptable reproducibility and precision with Clindamycin and the WalkAway® instrument.

Quality Control testing demonstrated acceptable results for Clindamycin.

DEPARTMENT OF HEALTH & HUMAN SERVICES





Food and Drug Administration 2098 Gaither Road Rockville MD 20850

Ms. Maureen Mende Regulatory Affairs Group Manager Dade Behring, Inc. 1584 Enterprise Boulevard West Sacramento, CA 95691-9972

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Re: 1

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Trade/Device Name: MicroScan MICroSTREP plus® Panel

Clindamycin (0.015 to 2 mcg/ml)

Regulation Number: 21 CFR § 866.1640

Regulation Name: Antimicrobial susceptibility test powder

Regulatory Class: II

Product Code: LRG, LTT Dated: August 28, 2006

Received: September 18, 2006

Dear Ms. Mende:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such additional controls. Existing major regulations affecting your device can be found in Title 21, Code of Federal Regulations (CFR), Parts 800 to 895. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Parts 801 and 809); and good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820).

This letter will allow you to begin marketing your device as described in your Section 510(k) premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus, permits your device to proceed to the market.

If you desire specific information about the application of labeling requirements to your device, or questions on the promotion and advertising of your device, please contact the Office of In Vitro Diagnostic Device Evaluation and Safety at (240)276-0450. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). Other general information on your responsibilities under the Act may be obtained from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 443-6597 or at its Internet address http://www.fda.gov/cdrh/dsma/dsmamain.html.

Sincerely yours,

Sally A. Hojvat, M.Sc., Ph.D.

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Director

Division of Microbiology Devices

Office of In Vitro Diagnostic Device

Evaluation and Safety

Center for Devices and Radiological Health

Enclosure

Indication for Use Statement

510(k) No.:	KO 6277 (To be assigned by FD	
Device Name:	MicroScan MICroSTREP <i>plus</i> ® Panel Clindamycin (0.015 to 2 mcg/ml)	
Intended Use	To determine bacterial antimicrobial agent susceptibility	
Indications for Use:	The MicroScan MICroSTREP plus® Panel is used to determine quantitative and/or qualitative antimicrobial agent susceptibility of colonies grown on solid media of aerobic streptococci, including Streptococcus pneumoniae. After inoculation, panels are incubated for 20 – 24 hours at 35°C +/- 1°C in a non-CO2 incubator, and read visually according to the Package Insert. Additionally, the panels may be incubated in and read by a MicroScan® WalkAway instrument.	
	This particular submission is for the addition of instrument read capability of the antimicrobial Clindamycin, at concentrations of 0.015 to 2 mcg/ml on the MicroScan MICroSTREP plus® Panel and the removal of the "Do not report" for <i>S. pneumoniae</i> .	
	The organisms which may be used for Clindamycin susceptibility testing on this panel are:	
•	Streptococci Streptococcus pn	eumoniae
Prescription Use X (Part 21 CFR 801 Subpart D)	AND/OR	Over-The-Counter Use(21 CFR 807 Subpart C)
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Concurrence of CDI	RH, Office of In Vitro Dia	
	Page 1 of1	Division Sign-Off
viii		Office of In Vitro Diagnostic Device Evaluation and Salety

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